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FILE 'HOME' ENTERED AT 14:56:20 ON 11 MAR 2005

=> file reg
COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
0.21 0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 14:56:29 ON 11 MAR 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 9 MAR 2005 HIGHEST RN 844817-50-1 DICTIONARY FILE UPDATES: 9 MAR 2005 HIGHEST RN 844817-50-1

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

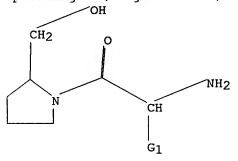
Please note that search-term pricing does apply when conducting SmartSELECT searches.

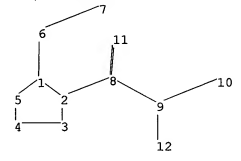
Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=>

Uploading C:\Program Files\Stnexp\Queries\10805624o.str





chain nodes :

6 7 8 9 10 11 12

ring nodes: 1 2 3 4 5

chain bonds : 1-6 2-8 6-7 8-9 8-11 9-10 9-12

ring bonds :

1-2 1-5 2-3 3-4 4-5

exact/norm bonds :

1-2 2-3 2-8 8-11 9-10 9-12

exact bonds :

1-5 1-6 3-4 4-5 6-7 8-9

isolated ring systems :

containing 1:

G1:C,H,Ak

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:CLASS 7:CLASS 8:CLASS 9:CLASS

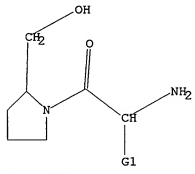
10:CLASS 11:CLASS 12:CLASS

L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR



G1 C, H, Ak

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 14:56:42 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 17596 TO ITERATE

5.7% PROCESSED 1000 ITERATIONS

0 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS:

343979 TO 359861

PROJECTED ANSWERS:

0 TO 0

PROJECTED ANSWERS:

0 SEA SSS SAM L1

=> s 11 full

L2

FULL SEARCH INITIATED 14:56:45 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 352611 TO ITERATE

95.0% PROCESSED 334980 ITERATIONS

15 ANSWERS

100.0% PROCESSED 352611 ITERATIONS

. 15 ANSWERS

SEARCH TIME: 00.00.26

=> file caplus
COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 161.33 161.54

FILE 'CAPLUS' ENTERED AT 14:57:15 ON 11 MAR 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

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FILE COVERS 1907 - 11 Mar 2005 VOL 142 ISS 12 FILE LAST UPDATED: 10 Mar 2005 (20050310/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 13 L4 15 L3

=> d ibib abs hitstr tot

L4 ANSWER 1 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
11TLE:
2004:435765 CAPLUS
141:140758
141:140758
Synthesis of D- and L-2,3-trans-3,4-cis-4,5-trans-3,4Dihydromy-5-hydromymethylproline and Tripeptides
Containing Then
Moreno-Vargas, Antonio J.; Robina, Inmaculada;
Petricci, Elena; Vogel, Pierre
Laboratorie de Glycochinie et de Synthese Asymetrique,
Swiss Federal Institute of Technology (EPFL),
Lausanne-Dorigny, CH-1015, Switz.
Journal of Organic Chemistry (2004), 69(13), 4487-4491
CODEM: JOCEAM; ISSN: 0022-3263
American Chemical Society
Journal CH-1015, SWITZ.
LANGUAGE:
CHEMISTORY TYPE:
LANGUAGE:
CASREACT 141:140758

111 ۲V

Enantiomerically pure (-)- and (+)-7-(tert-butoxycarbonyl)-5,6-exoisopropylidenedioxy-7-azabicyclo[2.2.1]heptan-2-ones, I and II, resp.,
were prepared I and II were converted into D- and L-2,3-trans-3,4-cis-4,5trans-N-(tert-butoxycarbonyl)-5-hydroxymethyl-3,4isopropylidenedioxyprolines, III and IV, resp. Applying the Boc and Fmoc
strategies of peptide synthesis, these compds were used to construct two
tripeptides. For example, III was incorporated into peptide synthesis to
give tripeptide V.
726192-28-59

RL: SPN (Synthetic preparation); PREP (Preparation) (asym. preparation of (dihydroxy)hydroxymethylproline and its incorporation

rporation into tripeptides) 726192-28-5 CAPLUS L-Valine, D-alanyl-(3S,4R,5R)-3,4-dihydroxy-5-(hydroxymethyl)-D-prolyl-,

L4 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
139:334930 CAPLUS
139:334930 CAPLUS
139:331666
Method for re-sensitizing vancomycin resistant bacteria using agents which selectively cleave a cell wall depsipeptide
Chiosis, Gabriela?
Chiosis, Gabriela? Boneca, Ivo G., Still, W. Clark
The Trustees of Columbia University in the City of New York, USA
PCT Int. Appl., 105 pp.
CODEN: PIXXVD2
DOCUMENT TYPE:
Fatent
LINGUAGE:
English

English

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WO 2003035098 20030501 WO 2002-US26975 20020823 Al

PRIORITY APPLN. INFO.:

OTHER SOURCE(s): MARPAT 138:331666

The present invention relates a method for re-sensitizing vancomycin resistant Gram-pos. bacteria in which resistance results from the conversion of an amide bond to an ester bond in the cell wall peptide precursors of the bacteria which comprises using an antibacterial amount of vancomycin or a homolog of vancomycin and an amount of an agent effective to selectively cleave the ester bond to thereby re-sensitize vancomycin resistant bacteria.

17 376643-20-89

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); TRU (Therapeutic use); BIOL (Biological study); PREF (Preparation); USES (Uses)

(Uses)
(re-sensitizing vancomycin resistant Gram-pos. bacteria using agents which selectively cleave ester bond of D-Ala-D-Lac cell wall depsipeptide)
376643-20-8 CAPLUS
2-Pyrrolidinemethanol, 1-(aminoacetyl)-, (2S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 1 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN phenylmethyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

518012-31-2

RR: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (re-sensitizing vancomycin resistant Gram-pos. bacteria using agents which selectively cleave ester bond of D-Ala-D-Lac cell wall

depsipeptide)
518012-31-2 CAPLUS
2-Pyrrolidinemethanol, 1-[(2S)-2-amino-1-oxopropyl]-, (2S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LA ANSWER 3 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 2001:643886 CAPLUS
DOCUMENT NUMBER: 136:2743
TITLE: Selective cleavage of D-Ala-D-Lac by small molecules:
re-sensitizing resistant bacteria to vancomycin
AUTHOR(S): Chicais, Gabrielar Boneca, Ivo G.
CORPORATE SOURCE: Department of Chemistry, Columbia University, Nev
York, NY, 10027, USA
SCIENCE (Washington, DC, United States) (2001),
293(5534), 1484-1487
CODEN: SCIEMS: 158N: 0036-8075
PUBLISHER: American Association for the Advancement of Science
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Pathogenic enterococci are becoming resistant to currently available
antibiotics, including vancomycin, the drug of last resort for Gram-pos,
infections. Enterococci pose a significant public health threat, not
least because of the risk of transferring vancomycin resistance to the
ubiquitous Staphylococcus aureus. Vancomycin resistance to the
ubiquitous Staphylococcus aureus. Vancomycin resistance to the
assembly and complementary chirality to the peptidoglycan termin that cannot bind
the antibiotic. Small mols. with well-oriented nucleophile-electrophile
assembly and complementary chirality to the peptidoglycan termin were
identified as catalytic and selective cleavers of the peptidoglycan
precursor depsiapeptide. These mols. were tested in combination with
vancomycin and were found to re-sensitize vancomycin-resistant bacteria to
the antibiotic.
IT 378643-19-5 378643-20-8
RL: BSU (Biological Study, unclassified); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(Resistive cleavage of b-Ala-D-Lac by small mols.: re-sensitizing
resistant bacteria to vancomycin)

NO 376643-19-5 CAPIUS

Absolute stereochemistry.

Absolute stereochemistry.

Same

376643-20-8 CAPLUS 2-Pyrrolidinemethanol, 1-(aminoacetyl)-, (2S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2000:539139 CAPLUS DOCUMENT NUMBER: 133:277734

DOCUMENT NUMBER: TITLE:

133:277734
The degradation of glycoproteins with lithium borohydride: isolation and analysis of O-glycopeptides with reduced C-terminal amino acid residue Arbatsky, N. P., Likhoshestov, L. M., Serebryakova, M. V., Brusov, O. S., Shibaev, V. N., Derevitskaya, V. A., Rochetkov, N. K. Zelinskii Institute of Organic Chemistry, Russian Academy of Sciences, Moscow, 117334, Russia Russian Journal of Bioorganic Chemistry (Translation of Bioorganicheskaya Khimiya) (2000), 26(1), 45-53 CODEN: RJECET; ISSN: 1068-1620
MAIK Nauka/Interperiodica
Journal AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

CODEN: RABGET: ASSN: AUDO-1000

MAIK Nauka/Interperiodica

DOCUMENT TYPE: Journal

LANGUAGE: English

AB By the example of fetuin and a blood-group-specific mucin from porcine
stomach, we showed that, under conditions of reductive degradation of
glycoproteins with LiBER-LIOH in 70% acuseous tert-Bu alc., the reduction and
cleavage of amide bonds occur much faster than the simultaneous

B-elimination of carbohydrate chains O-linked with Ser and Thr
residues of the peptide chain. The major degradation products containing

tha

O-linked glycans are the O-glycosylated derivs. of 2-aminopropane-1,3-diol and 2-aminobutane-1,3-diol (the products of reduction of glycosylated Ser

Thr) and the glycopeptides containing 2-4 amino acid residues with reduced C-terminal amino acid. Seventeen homogeneous O-glycopeptides were isolated from the fetuin degradation products by ion-exchange and reversed-phase EFIC. Their structures were determined by MADIO-TOF mass spectrometry and by analyses for amino acids, amino alcs., and carbohydrates. The application of the reaction for characterization of O-glycans and localization of O-glycosylation sites in O- and N.O-glycoproteins is discussed.
299187-67-4

IT 299197-67-4

RL: BPR (Biological process): BSU (Biological study, unclassified): BIOL (Biological study): PROC (Process)

(structure of fetuin degradation products obtained by reductive degradation

with LiEH4-LiOH in aqueous tert-Bu alc.)

RN 299197-67-4 CAPLUS

CN 2-Pytrolidinesethanol, 1-[(25,3B)-3-[[O-(N-acetyl-a-neuraminosyl)-(2-3)-0-P-0-galactopyranosyl-(1-3)-2-(acetyl-amino)-2-deoxy-a-0-galactopyranosyl)(xyl-2-amino-1-oxobutyl)-, (25)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L4 ANSVER 3 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

REFERENCE COUNT: 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSYER 5 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1997:640667 CAPLUS
DOCUMENT NUMBER: 127:318974

ITILE: and analogs as protein tyrosine kinase pp60c-src inhibitors and analogs as protein tyrosine kinase pp60c-src inhibitors
Altmann, Eva Novartis A.-G., Svitz., Altmann, Eva Novartis A.-G., Svitz., Altmann, Eva CODER: PIXXOZ

DOCUMENT TYPE: PATENT INDORBATION: German
FAMILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

VD 9734895 A1 19970925 WD 1997-EP1095 19970305

W: AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GB, EU, IL, IS, JP, RP, RR, LC, LK, LA, LT, LV, HG, MK, HN, MX, MO, NZ, PL, RO, SG, SI, SK, TR, TT, UM, US, UZ, VM, TO, AM, AZ, SY, KG, KZ, HD, RU, TJ, TH

RW: GEI, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, IU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GM, ML, MR, NE, SM, TD, TG

CA 2247739 AA 19970925 CA 1997-2249739 19970305

AU 9721534 A1 19971010 AU 1997-21534 19970305

AU 9721534 A1 19971010 DA 1997-21534 19970305

EP 888353 A1 19990107 EP 1997-914189 19970305

EP 888353 A1 19990512 CN 1997-193899 19970305

R: AT, BE, CH, UE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, FI

CN 1216544 A 19990512 CN 1997-193899 19970305

EN 9709443 A 19990810 BR 1997-9443 19970305

JP 2000506537 T2 20000530 JF 1997-533081 19970305

AT 244719 E 20037128 AT 1997-314189 19970305

AT 244719 E 20037128 AT 1997-314189 19970305

ES 2203793 T3 20040416 ES 1997-914189 19970305

ES 2203793 T3 20040416 ES 1997-914189 19970305

EN 50804199 A 19981105 NO 1998-4199 19970305

PRIORITY APPLIAL INFO: CH 1996-694 A 19960315

PRIORITY APPLIAL INFO: CH 1996-694 A 19970305

OTHER SOURCE(S): MARPAT 127:318974 APPLICATION NO. PATENT NO. KIND DATE CH 1996-694 WO 1997-EP1095 OTHER SOURCE(S):

Title compds. [I; R = R52(CH2)0-4; R1 = aryl; R2,R3 = H, halo, alkyl; R5 = H, alkyl, alkanyl, alkonycatbonyl, etc.; Z = (un)substituted pyrcolidine-1,2- or 1,3-diyl, -piperidine-1,2-, -1,3-, or -1,4-diyl) were

MARPAT 127:318974

L4 ANSWER 6 OF 15
ACCESSION NUMBER:
DOCUMENT NUMBER:
115:136560
115:136560
Synthesis and biological evaluation of
4-purinylpyrrolidine nucleosides
Peterson, Mark L., Vince, Robert
CORPORATE SOURCE:
CORPORATE SOURCE:
USA

Journal of Medicinal Chemistry (1991), 34(9), 2787-97 CODEN: JMCMAR; ISSN: 0022-2623 SOURCE:

Journal DOCUMENT TYPE: English LANGUAGE:

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The synthesis of several novel carbocyclic purine nucleosides which incorporate a nitrogen in place of carbon 3 of the cyclopentyl moiety are described. These analogs are derived from the key stereoches. defined intermediate M-(tert-butoxycatohonyl)—O-(id-methoxyphenyl)diphenyl)minethyl-trans-4-hydroxy-D-prolinol (I), which was accessible in 61.10 overall yield for a five-step sequence starting from cis-4-hydroxy-D-proline. The heterocyclic bases, 6-chloropurine and 2-amino-6-chloropurine, are efficiently introduced onto the pyrrolidine ring via a Mitsunobu-type coupling procedure with 7h3F and di-Et szodicarboxylate. Standard transformations and removal of protecting groups gave the cis-adenine, hypoxanthine, 2,6-diaminopurine, and guanine D-prolinol derivs. II (K = H, Y = NBZ, GH; K = NBZ, Y = HBZ, GH). In addition, a related sequence from trans-4-hydroxy-L-proline provided the enantiomeric L-prolinol guanine derivative The 6-(dimethylamino)purine analog, was coupled to N-(benzyloxycatbonyl)-p-methoxy-L-phenylalanine to provide, after deprotection, the novel purcmycin-like analog III. The analogs II and III were evaluated for antitumor and virucidal activity. These compols, failed to appreciably inhibit the growth of P388 mouse leukenia cells in vitro at concns. up to 100 µg/nL. In addition, they did not exhibit noticeable activity against the HIV or herpes simplex virus type 1 at concns. as high as 100 µM. The adenian analog, I (X = H, Y = NBZ) proved to be a substrate for adenosine deaminase and possessed an affinity for the enzyme only 501 leps than that of adenosine viru ha Ki = 85 µM.

135042-36-39
RE: SPN (Synthetic preparation): PREP (Preparation)
(preparation, antileukemic, and virucidal activity of)
135042-36-3 CAPUS
2-Pyrrolidinemethanol, 1-[2-amino-3-(4-amithoxyphenyl)-1-oxopropyl]-4-[6(dimethylamino)-9H-purin-9-yl]-, [2R-[1(5*), 2m, 4m]]- (9CI)
(CA INDEX NAME)

Absolute stereochemistry.

ANSYER 5 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) prepd. as protein tyrosine kinase pp60C-src inhibitors (no data). Thus, Ph0CCHEMHAC was cyclocondensed with CH2 (CM)2 and the product condensed with HC(DE1)3 and NH3 to give N-(3-cyano-4-pheny)-2-pyrcolyllformamidine which was cyclized to give, after deprotection, I (RI = Ph, RZ = R3 = H) (III) R = H) which was condensed with Me (ER, 4R)-1-tert-butoxycarbomyl-4-tosyloxypyrcolidine-2-carbomylate to give, after deprotection, II (R = (PR, 45)-1-2-choxycarbomyl-4-pyrcolidinyl).

197525-26-1P

197925-26-IP
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of 7-heterocyclypyrrolol(2,3-d]pyrimidines and analogs as protein tyrosine kinase pp60c-src inhibitors)
197525-26-I CAPLUS
2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxopentyl)-4-(4-amino-5-phenyl-TH-pyrolol(2,3-d]pyrimidin-7-yl)-, dihydrochloride,
[2R-[1(2S*,3S*),2m,48]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry

●2 HC1

ANSWER 6 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

L4 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
1978:152891 CAPLUS
1

CODEN: ABCHA6: ISSN: 0002-1369

DOCUMENT TYPE:

COCHMeNH2

2,4-Diamino-2,4-dideoxy-L-arabinose derivs, were prepared from benzyl 2-(benzylozycarbonyl) amino-2-deoxy-B-D-glucofuranoside by a series of known reactions. Among the compds, prepared is furancid prumycin I. 66167-01-9.

RL: RCT (Reactant): SPM (Synthetic preparation): PREP (Preparation): PACT (Reactant or reagent)
(preparation and catalytic hydrogenolysis of)
66167-01-9 CAPLUS
Carbamic acid, [1-(2-mino-1-oxopropyl)-2,4-dihydroxy-5-(hydroxymethyl)-3-pyrcoldinyl-, phenylmethyl ester, (2R-[1(R*),2x,3x,4\$,5])

Absolute stereochemistry.

ΙT

66167-02-0P
RL: SPM (Synthetic preparation), PREP (Preparation)
(preparation of)
66167-02-0 CAPIUS
2,4-Pyrcolidinediol, 3-amino-1-(2-amino-1-oxopropyl)-5-(hydroxymethyl)-,
dihydrochloride, [2R-[1(R*),2m,3m,4P,5m]]- (9CI)
(CA INDEE NAME)

L4 ANSWER 8 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1975:459253 CAPLUS

BOCKNEWN NUMBER: 31:59253 CAPLUS

B3:59253 CAPLUS

B4:59253 CAPLUS

B3:59253 CAPLUS

B3:59253 CAPLUS

B3:59253 CAPLUS

B4:59253 CAPLUS

B

DOCUMENT TYPE: JOURNAL LANGUAGE: Brilish Alburghage Printed CA Issue.

AB The mass spectrum of actinonin (1) was interpreted by comparison with the fragmentation of the model compds. II-V. The structure of I, except for spectrum. the pos--spectrum.

IT 54124-50-6
Ri PRP (Properties)
(mass spectrum of)
RS 54124-60-6 CAPUS
CN 2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxobutyl)-, [S-(R*,R*)](9CI) (CA INDEX NAME)

L4 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

Absolute stereochemistry.

●2 HC1

L4 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1975:459252 CAPLUS
DOCUMENT NUMBER: 83:59252 APLUS
83:59252 Antibiotic actinonin. VI. Synthesis of structural analogs of actinonin by dicyclohexylcarbodiimide coupling reactions
Bevlin, John F., Ollis, W. David; Thorpe, John E., Wright, Derek E.
CORPORATE SOURCE: Dep. Chem., Univ. Sheffield, Sheffield, UK Journal of the Chemical Society, Perkin Transactions 1: Organic and Bio-Organic Chemistry (1972-1999) (1975), (9), 848-51 CODEN: JCPRB4; ISSN: 0300-922X

DOCUMENT TYPE:

UAGE: Journal
For diagram(s), see printed CA Issue.
Coupling of anino amides with monoesters of dicarboxylic acids with
dicyclohexylcarbodismide in CHZC12 gave dicarbamcyl esters, which with
MedH-MHZOH gave the corresponding hydroxamic acids, analogs of actinonin.
E.g., DL-valylmorpholine with HOZCCH[(CH2)4Me]COZEt gave the ester I,
which gave the hydroxamic acid II.
54124-60-6
RL: RCT (Reacters).

Selze-Bu-B RL: RCT (Reactant); RACT (Reactant or reagent) (coupling reaction with dicarboxylic acid monoesters) 54124-60-6 CAPLUS

2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxobutyl)-, [S-(R*,R*)]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 54124-60-69
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and reaction with methanolic hydroxylamine)
54124-60-6 CAPUS
2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxobutyl)-, [S-{R*,R*}](SCI) (CA INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1975:459248 CAPLUS
83:59248
Antibiotic actinonin. II. Total synthesis of actinonin and structural analogs by the isomaleimide method
AUTHOR(S):
Anderson, Nicholas H.; Ollis, W. David; Thorpe, John E.; Ward, A. David
Dep. Chem., Univ. Sheffield, Sheffield, UK
Journal of the Chemical Society, Perkin Transactions 1: Organic and Bio-Organic Chemistry (1972-1999)
(1975), (9), 825-30
CODEM: JORDA! JORDA! ISSN: 0300-922X
JOURNAL SHEET STREET S

DOCUMENT TYPE: Journal
LANGUAGE: English
GI For diagram(s), see printed CA Issue.
AB Valylprolinol with the isomaleimide I gave O-benzyldidehydroactinonin (II)
which on hydrogenation gave actinonin (III). Analogs IV-VI were prepared
similarly from alanylpyrrolidine, valylpyrrolidine, and valylprolinol,
reso.

resp. 54124-60-6P

54124-60-69
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and reaction with isomaleimide derivative)
54124-60-6 CAPLUS
2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxobutyl)-, [S-(R*,R*)](9CI) (CA INDEX NAME)

Absolute stereochemistry.

RL: SPN (Synthetic preparation); PREP (Preparation)

L4 ANSVER 10 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1975:459251 CAPLUS

S3:59251

TITLE: Antibiotic actinonin. V. Synthesis of structural analogs of actinonin by the anhydride-ester method analogs of actinonin by the anhydride-ester method analogs of actinonin by the anhydride-ester method by the anhydride-ester method analogs of actinonin by the anhydride-ester method to the component of the

Absolute stereochemistry.

L4 ANSWER 12 OF 15
ACCESSION NUMBER:
DOCUMENT NUMBER:
1975:459247 CAPLUS
S1:59247 Antibiotic actinonin. I. Constitution of actinonin.
Natural hydroxamic acid with antibiotic activity
Gordon, James J., Devlin, John P.; East, Anthony J.;
Ollis, W. David: Sutherland, Ian O.; Wright, Derek E.;
Ninet, Leon
CORPORATE SOURCE:
Antibiot. Res. Stat., Med. Res. Counc., Clevedon, UK
JOURNAI of the Chemical Society, Perkin Transactions
1: Organic and Bio-Organic Chemistry (1972-1999)
(1975), (9), 819-25
CODEN: JOURNAI
LANGUAGE:
The structure of actinonin (1), isolated from Streptomyces roseopallidus,
was determined by degradation to its constituent residues, L-prolinol,
Parentylaucrics acid, and bydrowylanine and from spectral data

ne, D-pentylauccinic acid, and hydroxylamine and from apectral data. 56439-51-1P

IT

56439-51-1P
RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)
56439-51-1 CAPUS
2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxobutyl)-, [S-(R*,R*)]-,
compd. with 2,4,6-trinitrophenol (1:1) (9CI) (CA INDEX NAME)

CH 1

CRN 54124-60-6 CMF C10 H20 N2 O2

Absolute stereochemistry.

2

CRN 88-89-1 CMF C6 H3 N3 O7

L4 ANSVER 13 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1974:535864 CAPLUS
TITLE: 81:135864
TOTAL SYNTHEMS: 6, 1135864
TOTAL SYNTHEMS: 81:135864
TOTAL SYNTHEMS: 6, 1011:2, W. David; Thorpe, John E.; Ward, A. David
CORPORATE SOURCE: Dept. Communications (1974), (11), 420-1
COUNTRY TYPE: Journal of the Chemical Society, Chemical Communications (1974), (11), 420-1
COUDEN; JOCCAT: ISSN: 0022-4936
JOURNAL Explish
OF of diagram(s), see printed CA Issue.
AB A regioselective and streeoselective synthesis of actinonin (I) from condensation of pentylnaleic anhydride with PhcHZONHZ was described.
IT 54124-60-6
RE: RCT (Reactant); RACT (Reactant or reagent)
(addition reaction with isomaleimide)
N 56124-60-6 CAPUS
CN 2-Pyrrolidinemethanol, 1-(2-amino-3-methyl-1-oxobutyl)-, [5-(R*,R*)](9CI) (CA INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1966: 482599 CAPLUS
DOCUMENT NUMBER: 65:82599
ORIGINAL REFERENCE NO.: 65:15497c-d
FITLE: Partial acid hydrolysis of \(\gamma\)-keratose
AUTHOR(S): Asquith, R. S.; Shaw, T.
COMPORATE SOUNCE: Bradford Inst. Tech., Bradford, UK
SOURCE: Journal
LANGUACE: Legilsh
AB \(\gamma\)-Keratose was hydrolyzed 192 hrs. in SN HCl at 37° to obtain
a hydrolyzate in which, based on amino N determination, the average peptide
chain
length was 2 amino acid residues. The partial hydrolyzate was
fractionated by ion exchange chromatography, two dimensional paper
chromatography, and/lor) high voltage paper electrophoresis. Fifteen diamino acid residues were identified and other peptides containing up to 5
amino acid residues also were found. Cysteylcysteic acid was shown to be
present.

amino acid residues also were found. Cysteylcysteic acid was shown present.
7754-78-1, p-Toluenesulfonamide, N-[[4-amino-4-[[2-(hydroxymethyl)-1-pyrrolidinyl]carbonyl]butyl]amidino][preparation of)
7754-78-1 CAPIUS
Pyrrolidine, 2-(hydroxymethyl)-1-[NS-[(p-tolylsulfonyl)amidino]-Lornithyl]-, L- (8CI) (CA INDEX NAME)

PAGE 1-A

L4 ANSVER 14 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
DSCUMENT NUMBER:
B8:108480 Unconventional nucleotide analogs. XI. Synthesis of a nonsaccharidal analog of purconycin
AUTHOR(S): Raspersen, Frans M.; Bieraugel, Hans; Pandit, Upendra K.

AUTHOR(S):

a nonsaccharidal analog of purconycin
Raspersen, Frans M.; Bieraugel, Hans; Pandit, Upendra K.

CORPORATE SOURCE:
Org. Chen. Lab., Univ. Amsterdam, Amsterdam, Neth.
BOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
LINKGUAGE:
English
GI For diagram(s), see printed CA Issue.
AB The title purcoycin analog (I), of interest because of analogy to nucleo-peptide models, is prepared Thus, (-)-4-hydroxy-1-proline was converted to II which on treatment with 5-amino-4,6-dichloropyrtaidine followed by ring closure ([Eto] 3CH] gave III (R - Cl., Rl = tosyl).
Reaction of this with Mc2MH and detosylation gave III (R - MMe2, Rl = H).
Coupling of this with Cbz N-protected 4-MeOCGH4CH2CH(NH2)-CO2H gave, after removal of the Cbz group, I.

II 51950-02-8P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
RN 51950-02-8P
RS: Synthetic preparation); PREP (Preparation)
(preparation of)
RN 1970-014 (Preparation) (PREP) (Preparation)
(preparation of)
RN 1970-014 (Preparation) (PREP) (Preparation)
(CA INDEX NAME)
Absolute stereochemistry.

Absolute stereochemistry.

L4 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

PAGE 2-A